PCT

WORLD INTELLECTUAL PROPERTY ORGANIZATION International Bureau



INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

(51) International Patent Classification 7:
 B01J 23/02, 23/58, 37/02, B01D 53/94

(11) International Publication Number:

WO 00/50168

A1 |

(43) International Publication Date:

31 August 2000 (31.08.00)

(21) International Application Number:

PCT/GB00/00571

(22) International Filing Date:

17 February 2000 (17.02.00)

02.001

(30) Priority Data:

9904269.9

24 February 1999 (24.02.99)

GB

(71) Applicant (for all designated States except US): EUROPEAN COMMUNITY represented by THE COMMISSION OF THE EUROPEAN COMMUNITIES [LU/LU]; Rue Alcide De Gasperi, L-2920 Luxembourg (LU).

(72) Inventors; and

- (75) Inventors/Applicants (for US only): PICKERING, Stephen [GB/NL]; Noordlaan 14, NL-1861 GN Bergen (NL). DJURICIC, Boro [NL/NL]; Mozartlaan 10, NL-1817 GS Alkmaar (NL). MAGNE, Jean-François [FR/CH]; Am Aawasser 10, CH-6056 Kägiswil (CH).
- (74) Agent: BOULT WADE TENNANT; Verulam Gardens, 70 Gray's Inn Road, London WC1X 8BT (GB).

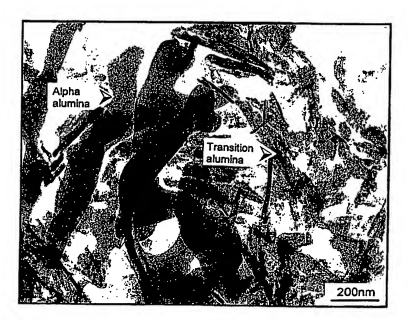
(81) Designated States: CA, JP, NO, US, European patent (AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE).

Published

With international search report.

Before the expiration of the time limit for amending the claims and to be republished in the event of the receipt of amendments.

(54) Title: STABILIZATION OF TRANSITION ALUMINA



(57) Abstract

The present invention relates to a process for the production of a precursor material for use in the preparation of a stabilized alumina catalyst support material, which process comprises the steps of: (a) providing a mixture comprising aluminium and barium containing compounds; and (b) heating the mixture under conditions to form particles of boehmite at least partially coated with a layer comprising barium and/or a compound of barium.

FOR THE PURPOSES OF INFORMATION ONLY

Codes used to identify States party to the PCT on the front pages of pamphlets publishing international applications under the PCT.

AL	Albania	ES	Spain .	LS	Lesotho	SI	Slovenia	
AM	Armenia	FI	Finland	LT	Lithuania	SK	Slovakia	
AT	Austria	FR	France	LU	Luxembourg	SN		
ΑU	Australia	GA	Gabon	LV	Latvia	SZ	Senegal Swaziland	
AZ	Azerbaijan	GB	United Kingdom	MC	Monaco	TD	Chad	
BA	Bosnia and Herzegovina	GE	Georgia	MD	Republic of Moldova	TG		
BB	Barbados	GH	Ghana	MG	Madagascar	TJ	Togo	
BE	Belgium	GN	Guinea	MK	The former Yugoslav	TM	Tajikistan Turkmenistan	
BF	Burkina Faso	GR	Greece		Republic of Macedonia	TR		
BG	Bulgaria	HU	Hungary	ML	Mali	TT	Turkey	
BJ	Benin	1E	Ireland	MN	Mongolia	UA	Trinidad and Tobago Ukraine	
BR	Brazil	IL	Israel	MR	Mauritania	UG		
BY	Belarus	IS	Iceland	MW	Malawi	US	Uganda	
CA	Canada	IT	Italy	MX	Mexico	UZ	United States of America	
CF	Central African Republic	JP	Japan	NE	Niger	VN	Uzbekistan	
CG	Congo	KE	Kenya	NL	Netherlands	YU	Vict Nam	
CH	Switzerland	KG	Kyrgyzstan	NO	Norway	ZW	Yugoslavia	
CI	Côte d'Ivoire	KP	Democratic People's	NZ	New Zealand	ZW	Zimbabwe	
CM	Cameroon		Republic of Korea	PL	Poland			
CN	China	KR	Republic of Korea	PT	Portugal			
CU	Cuba	KZ	Kazakstan	RO	Romania			
CZ	Czech Republic	LC	Saint Lucia	RU	Russian Federation			
DE	Germany	LI	Liechtenstein	SD	Sudan			
DK	Denmark	LK	Sri Lanka	SE	Sweden			
EE	Estonia	LR	Liberia	SG	Singapore			
				50	2 mg afronc			

WO 00/50168 PCT/GB00/00571

Stabilization of Transition Alumina

The present invention relates to the general field of catalysts, such as automobile exhaust catalysts, and, in particular, to a stabilized transition alumina catalyst support material.

Rapid developments in combustion technology coupled with the increasing demands for reduced levels 10 of pollutants emitted by combustion processes has created a demand for new catalyst materials. particular, new catalysts are needed for temperatures above 1000°C, not only for the removal of pollutants, for example to clean up automobile exhausts, but also 15 for catalytic combustion, for example in heaters, boilers, and gas turbines. The catalytic combustion of fuels offers the advantages of a more uniform combustion temperature and a more complete combustion of the fuel. The result is a higher thermal 20 efficiency, together with less nitrogen oxide formation through the prevention of localised high temperature areas.

automobile exhaust catalysts are used to reduce
automobile exhaust emissions by converting CO,
unburned hydrocarbons and NO, in the exhaust gas to an
environmentally less harmful mixture comprising CO,
H2O and N2. The catalysts typically comprise a metal
or ceramic monolithic carrier, such as Cordierite
(2MgO.2Al2O3.5SiO2), coated with a high surface area
catalyst support material. Catalytically active noble
metal particles such as, for example, Pt, Pd and/or
Rh, and other additives, are dispersed on and within
the support material.

5

WO 00/50168 PCT/GB00/00571

- 2 -

The catalytically active particles need to be held in the exhaust gas stream on a support material which has a high specific surface area to maximize their exposure to the gas stream and thereby maximize the catalytic efficiency. Gamma-alumina and the closely related transition aluminas theta and delta are commonly used as support materials because of their high specific surface areas. In particular, gamma-alumina is unusual among ceramic materials in that it has a very high specific surface area, for example exceeding 100 m^2/g , compared with the more typical values of from about 1 to 5 m^2/g for other ceramic materials. However, gamma and the other transition aluminas are metastable phases, which are subject to thermal degradation and undergo a phase transition at temperatures exceeding about 1000°C to the thermodynamically more stable alpha-alumina (see Figures 1 and 9B) with the corollary of a significant reduction in the specific surface area; typically from about 100 m^2/g to about 1 to 2 m^2/g . This results in a corresponding reduction in catalytic activity. The transformation sequence for boehmite is gamma \rightarrow delta \rightarrow theta \rightarrow alpha. Gamma, delta and theta alumina all have a cubic oxygen sub-lattice and the only difference between them is in the progressive increase in ordering of aluminium atoms on the cation deficient aluminium sub-lattice. In contrast, alpha-alumina has a hexagonal oxygen sub-lattice and its formation involves a total restructuring of the lattice.

30

35

5

10

15

20

25

Thermal degradation of transition alumina is a kinetic effect which accelerates rapidly with increasing temperature. Whereas at 600°C it is too slow to be significant within the lifetime of a catalytic converter, at 1100°C it may be complete

10

15

within 1 hour. Catalytic converters are operated at the upper limit of their stability and catalyst support materials with improved resistance to thermal degradation would allow higher operating temperatures and a longer service life.

Automotive exhaust gases from petrol engines may contain up to 3% CO and 100 ppm unburned hydrocarbons which react exothermically in the catalytic converter with NO_x to yield an environmentally less harmful mixture of gases, namely CO₂, H₂O and N₂. Peak operation temperatures of from about 700°C to about 800°C are frequently achieved. The current trend to reduce CO₂ emissions has resulted in increased use of diesel engines with lower fuel consumption and hence lower CO₂ emissions. However, diesel engines operate at higher exhaust temperatures; approximately 800°C is normal with peak temperatures of up to about 1300°C.

20 The conventional solution to the thermal degradation of gamma and the other transition aluminas is the use additives, such as, for example, Zr, Ba, Ce, La, Si, either singly or in combination. US-4,624,940 relates to a catalyst for the oxidation of carbon monoxide and unburned hydrocarbons in an 25 exhaust stream. The catalyst comprises support particles of stabilized gamma-alumina having palladium dispersed thereon and can remain effective after exposure to temperatures greater than 1000°C. gamma-alumina is stabilized by either (i) 30 incorporation of a combination of lanthana and baria into the alumina particles, or (ii) impregnation of the alumina particles with an aqueous dispersion of a high molecular weight polyorganosiloxane followed by heating to a temperature which is effective to 35

WO 00/50168

5

10

15

. 35

decompose the polyorganosiloxane. The additives are introduced by soaking an alumina powder or a solid alumina precursor in a solution containing the additive followed by drying and calcining at a temperature sufficient to effect stabilization. An alternative but less widely used method of introducing additives is by a coprecipitation process as described in US-4,056,489. In this manner an intimate admixture of alumina and the stabilizing additives is formed, which is then calcined.

The present inventors have now devised a process whereby transition alumina can be further stabilized against thermal degradation to alpha-alumina. This results in a catalyst which is effective at higher operating temperatures and for longer times compared with the prior art catalysts.

Accordingly, in a first aspect the present
invention provides a process for the production of a
precursor material for use in the preparation of a
stabilized alumina catalyst support material, which
process comprises the steps of:

- 25 (a) providing a mixture comprising an aluminium containing compound and at least one additive selected from barium, zirconium, cerium, lanthanum, calcium and silicon containing compounds; and
 - (b) heating the mixture under conditions to form particles of boehmite at least partially coated with a layer comprising at least one of barium, zirconium, cerium, lanthanum, calcium and silicon and/or at least one compound containing barium,

zirconium, cerium, lanthanum, calcium and/or silicon.

Boehmite is a mineral form of aluminium monohydroxide (AlO.OH).

The additive is preferably a barium containing compound. The layer formed on the particles of boehmite preferably comprises barium carbonate.

An important advantage of the process according to the present invention is that the particles of boehmite can be substantially coated completely with the said layer.

The mixture in step (a) is preferably a substantially homogenous mixture. The substantially homogenous mixture may be formed by any process which results in an intimate admixture. For example, by forming a slurry of boehmite and/or gamma alumina with a water-soluble compound of barium, such as barium nitrate, or, alternatively, by adding a barium containing solution to a gel comprising an aluminium containing compound.

20

25

30

35

5

10

15

A particularly preferred process for forming the substantially homogenous mixture is by homogeneous precipitation, ideally homogeneous precipitation of water-soluble aluminium and barium containing compounds, for example salts thereof. Suitable aluminum compounds include aluminium nitrate, aluminium sulphate and/or aluminium chloride, preferably aluminium chloride hexahydrate (AlCl₃·6H₂O). Suitable barium compounds include barium nitrate, barium sulphate and/or barium chloride, preferably barium chloride dihydrate (BaCl₂·2H₂O).

Precipitation from aqueous solution is widely used for producing ceramic oxide powders.

Precipitation occurs by adding a precipitating ligand

(anion) to a solution containing cations of the appropriate metal. If the precipitating ligand is added directly by simply pouring one solution into another then there is little control of the chemistry during precipitation because of the large and 5 inhomogeneous gradients in solution concentration. Better control of chemical and morphological characteristics can be achieved if the precipitating ligands are generated "in situ", simultaneously and uniformly throughout the solution. This is know as a 10 "homogeneous" precipitation process. In a preferred aspect of the present invention the mixture in step (a) is formed by homogeneous precipitation in which a base is generated within a metal salt solution through thermal decomposition of a suitable water-soluble 15 reagent. Suitable water-soluble reagents include any compound that decomposes on heating in aqueous solution to release NH3 groups, thereby forming ammonium hydroxide which increases the pH of the solution. Examples include urea and amines, such as 20 hexamethylene tetramine. Suitable compounds of aluminium include aluminium nitrate, aluminium sulphate and/or aluminium chloride, preferably aluminium chloride hexahydrate (AlCl₃·6H₂O). compounds of barium include barium nitrate, barium sulphate and/or barium chloride, preferably barium chloride dihydrate (BaCl₂·2H₂O). A small amount of a dispersant, for example polyvinylpyrrolidone, may also be added to the metal salt solution. The dispersant improves the uniformity of the precipitation product by reducing the tendency of the precipitate particles to stick together. Additionally, diffusion of ionic species through an adsorbed dispersant-layer may be the rate limiting step to further particle growth.

30

25

15

30

35

The mixture comprising aluminium and barium containing compounds will generally be in the form of a gel when produced by the homogeneous precipitation process described above, although the mixture can take other forms including, for example, hydroxides, 5 molecular and ionic complex species or compounds. gel is a three-dimensional network of colloidal particles of mainly amorphous aluminium hydroxide containing a large amount of water (for example approximately 170 g of wet gel yields approximately 7 g of dried powder) and in which the barium additive is trapped. The consistency of the gel is generally between that of a thick paste and a jelly, and it easily breaks up when stirred in an excess of water and slowly settles to the bottom again.

The mixture comprising aluminum and barium containing compounds may also be formed by adding a water-soluble barium compound, for example barium nitrate, barium sulphate and/or barium chloride, to a 20 slurry of, for example, boehmite in water. Advantageously, an organic acid, such as a carboxylic acid is added to the mixture. A preferred carboxylic acid is oxalic acid. In a particularly preferred aspect, a solution comprising barium nitrate and 25 oxalic acid is added to the slurry of boehmite in water and the mixture is then heated under conditions to produce particles of boehmite at least partially coated with a layer comprising barium and/or a barium compound.

The heating in step (b) is preferably achieved by a hydrothermal treatment, which is typically carried out on an aqueous solution of the mixture of aluminium and barium containing compounds in a pressure vessel,

such as an autoclave, at a temperature above the normal boiling point and under applied or autogenous pressure to substantially prevent boiling. hydrothermal treatment will generally be performed at a temperature typically in the range of from 90 to 5 300°C, more typically from 100 to 220°C. It will be appreciated that the pressure required to substantially prevent boiling will depend on the applied temperature. Typically, the hydrothermal treatment is performed at a pressure in the range of 10 from 1 to 150 bar, more typically from 5 to 50 bar, still more typically from 10 to 40 bar. hydrothermal treatment is typically carried out for a time in the range of from 30 minutes to 25 hours, more 15 typically from 1 to 10 hours.

The heating in step (b) may also be achieved by heating at a temperature in the range of from ambient temperature to boiling temperature. This may readily be achieved by a reflux operation, which may be carried out, for example, at atmospheric pressure and a temperature in the range of from 80 to 120°C, preferably approximately 100°C.

In one aspect of the present invention, the process of forming coated boehmite particles from a solution comprising aluminium and barium containing compounds can be seen to involve sequential precipitation: first, the formation of an aluminium hydroxide gel; second the precipitation of barium and/or a barium compound (probably as a carbonate) within the gel at a higher pH to give a uniform mixture; third crystallization of boehmite particles; and fourth precipitation or re-precipitation of barium and/or a barium compound (probably as a carbonate) on

the surface of the boehmite particles.

The product of the process according to the first aspect of the present invention may subsequently be dried and then calcined. Drying and calcination will generally be carried out in air or other oxidizing atmosphere.

In a second aspect, the present invention

provides a process for the preparation of a stabilized alumina catalyst support material, which process comprises the steps of:

- (i) providing a precursor material comprising
 particles of boehmite at least partially coated
 with a layer comprising barium and/or a compound
 of barium; and
- (ii) heating the coated particles of boehmite to a temperature at which at least some of the boehmite transforms to gamma-alumina.

In the second aspect, heating is typically carried out at a temperature in the range of from 500 to 1000°C. The process may also include an additional step of further heating whereby at least some of the gamma-alumina transforms to theta and/or delta-alumina. The further heating is typically carried out at a temperature in the range of from 1000 to 1400°C.

Heating will generally be carried out in air or other oxidizing atmosphere.

The precursor material comprising particles of boehmite may be unsupported or may be provided in the form of a washcoat on a catalyst carrier substrate,

15

20

such as a ceramic honeycomb.

The gamma-alumina particles produced according to the second aspect of the present invention may be provided in the form of a dry powder which can be redispersed in an appropriate liquid, such as water, to form a slurry or washcoat.

The precursor material may be produced according to a process as herein described with reference to the first aspect of the invention.

In a third aspect, the present invention provides a process for the preparation of a catalyst, such as an automotive catalytic converter, which process comprises the steps of dispersing a precursor mixture comprising particles of boehmite at least partially coated with a layer comprising barium and/or a compound of barium on a metal or ceramic monolithic substrate, followed by heating the precursor mixture to a temperature at which at least some of the boehmite transforms to gamma-alumina.

In the third aspect, heating is typically carried out at a temperature in the range of from 500 to 1000°C. The process may also include an additional step of further heating whereby at least some of the gamma-alumina transforms to delta and/or theta-alumina. The further heating is typically carried out at a temperature in the range of from 1000 to 1400°C. Again, heating will generally be carried out in air or other oxidizing atmosphere.

The mixture can be dispersed on to metal or ceramic monolithic substrate in a conventional manner,

35

for example by dipping the substrate one or more times into an aqueous slurry (washcoat) of the coated boehmite powder, followed by drying.

In the second and third aspects of the present invention, the particles of gamma-alumina, theta, delta-alumina and/or boehmite will typically be at least partially coated with a layer comprising barium and/or a compound of barium. The layer will typically comprise one or more of barium carbonate, barium oxide and/or barium aluminate depending on the heat-treatment. For gamma-alumina particles, the layer will generally comprise barium oxide as the major constituent, whilst for theta and delta-alumina particles, the layer will generally comprise one or more barium aluminates as the major constituent(s).

In a fourth aspect, the present invention provides a process for the preparation of a catalyst, such as an automotive catalytic converter, which process comprises the step of dispersing a stabilized alumina catalyst support material as herein described on a metal or ceramic monolithic substrate.

Again, the stabilized alumina catalyst support material may be dispersed on to the metal or ceramic monolithic substrate in a conventional manner, for example by dipping the substrate one or more times into an aqueous slurry (washcoat) of stabilized alumina powder, followed by drying.

In a fifth aspect, the present invention provides a precursor material for use in the preparation of catalyst support material, which precursor material comprises particles of boehmite and/or transition alumina at least partially coated with a layer comprising barium and/or a compound of barium.

Each particle of boehmite and/or transition alumina preferably has a substantially uniform surface 5 layer comprising barium and/or a compound of barium. The layer will typically comprise one or more of barium carbonate, barium oxide and/or barium aluminate depending on the heat-treatment used to process the particles. For boehmite, the layer will generally 10 comprise barium carbonate as the major constituent; for gamma-alumina particles, the layer will generally comprise barium oxide as the major constituent; for theta and delta-alumina particles, the layer will generally comprise one or more barium aluminates as 15 the major constituent(s).

The size of the particles may range from 10 nm to 5 μm, but the particles typically have an average particle size in the range of from 100 to 500 nm. In general the particles take the form of thin prismatic crystals with an average length to width ratio typically of about 3:1. The average length to thickness ratio is typically in the range of from 10:1 to 20:1. The crystal sizes are for the longest dimension. The surface layer comprising barium and/or a compound of barium will typically be up to 5 nm in thickness, more typically up to 3 nm in thickness.

The precursor material may provided in the form of an aqueous slurry, a non-aqueous slurry or a powder.

In a sixth aspect, the present invention provides a stabilized alumina catalyst support material which

25

35

comprises gamma, theta and/or delta-alumina, and which withstands substantial degradation to alpha-alumina after exposure to a temperature of approximately 1400°C for 1 hour when heated in, for example, air or other oxidizing atmosphere.

In this aspect, the stabilized alumina catalyst support material preferably has a specific surface area of at least 20 m²/g, more preferably at least 25 m²/g, still more preferably at least 31 m²/g, after exposure to a temperature of 1400°C for 1 hour. The specific surface area is measured using the standard BET method.

In a seventh aspect, the present invention provides a catalyst, such as an automotive catalytic converter, which comprises a metal or ceramic monolithic substrate having dispersed thereon a stabilized alumina catalyst support material as herein described.

It will be appreciated that one or more catalytically active noble metal particles, for example Pt, Pd and/or Rh, optionally together with other additives, will typically be dispersed on the support material.

In the third, fourth and seventh aspects the catalyst may be a 3-way automotive catalytic converter.

The monolithic substrate can be formed from any of the conventional ceramic materials in the art including, for example, one or more of cordierite, cordierite-alpha alumina, silicon nitride, silicon

carbide, mullite, zirconia-mullite, spodumene, alumina-silica-magnesia and/or zirconium silicate. Examples of suitable metal substrates include austenitic stainless steel and high aluminium ferritic stainless steel.

It will be appreciated that the stabilized alumina according to the present invention may be used in conjunction with other components conventionally used in the field of catalysis, such as, for example: catalytic promoters, including Ni, Co, Mn, Cr₂O₃, TiO₂, ZrO₂ and/or CeO₂; washcoat stabilization oxides, including cordierite, mullite and/or magnesium aluminium titanate; additional stabilization

15 additives, including Zr, Ce, La, Ca, Cu, Th and/or Si.

The present invention will now be described further, by way of example, with reference to the following drawings, in which:

20

5

Figure 1 is a TEM micrograph of alumina without additive showing the transformation of transition-alumina to alpha-alumina with a reduction in specific surface area;

25

30

35

Figure 2 is a TEM micrograph of a hydrothermally treated material showing large alpha-alumina crystals and nanoparticles of barium aluminate. The material was formed by heating a hydrothermally treated alumina-baria precursor to 1400°C for 1 hour in argon;

Figure 3 is a TEM micrograph showing a porous boehmite crystal pseudomorph consisting of gamma-alumina coated with a barium oxide layer. The

20

25

material was formed by heating a hydrothermallytreated precipitate of an alumina-baria precursor to 1000°C for 1 hour;

Figures 4 shows TEM micrographs of boehmite pseudomorphs consisting of theta-alumina particles exhibiting a twinned structure and some residual porosity. The material was formed by heating a hydrothermally-treated precipitate of alumina-baria precursor to 1400°C for 1 hour;

Figure 5 A and B are TEM micrographs showing a comparison between boehmite produced by a hydrothermal treatment of a precipitate of an alumina-baria precursor and theta-alumina formed by heating the same boehmite: (A) boehmite produced by a hydrothermal treatment of a precipitate of an alumina-baria precursor, and (B) theta-alumina formed by heating the boehmite to 1400°C for 1 hour;

Figure 6 is a flow chart showing a synthesis procedure for making alumina-baria precursor gels;

Figure 7 is a DTA analysis of an autoclave treated alumina-baria precursor gel for various heating rates.

Figures 8 A-D are TEM micrographs of an oven-dried alumina-baria precursor gel: (A) and (B) microcrystalline bayerite before calcination, bright and dark filed respectively, (C) and (D) gamma-alumina with crystals of β_{II} after 1 hour at 1000°C, bright and dark field respectively;

10

15

30

35

Figures 9 A-F are TEM micrographs of a microwave treated alumina-baria precursor gel: (A) morphology after microwave drying, (B) large alpha-alumina crystal growing in matrix of theta-alumina after 1 hour at 1200°C in air, (C) and (D) nano-particles of β_i inside alpha-alumina crystals after 1 hour at 1400°C in air, (E) and (F) nano-particles of β_i inside and on the surface of alpha-alumina crystals after 1 hour at 1400°C in argon;

Figure 10 A-C are TEM micrographs of an autoclave treated alumina-baria precursor gel: (A) well-developed boehmite crystals after hydrothermal treatment of 4 hours at 210°C, (B) porous boehmite pseudomorphs after 1 hour at 1000°C, and (C) detail showing size, shape and orientation of pores;

Figures 11 A-D are TEM micrographs of an autoclave treated alumina-baria precursor gel after calcination for 1 hour at 1400°C in air:

(A) boehmite pseudomorphs comprising theta-alumina, (B) finely twinned theta-alumina with equiaxed pores, (C) and (D) details of twinned theta-alumina structure showing twin lamella as narrow as 5 nm; and

Figures 12 A and B are TEM micrographs of an autoclave treated alumina-baria precursor gel after calcination for 1 hour at 1400°C in argon:

(A) nano-particles of BaAl₁₂O₁₉ in a matrix of theta-alumina transformed extensively to alpha-alumina, (B) detail showing BaAl₁₂O₁₉ particles predominantly on the surface of the alpha-

10

35

alumina.

The present inventors have investigated a number of ways of introducing barium additives to gamma-alumina and have found that they all yield similar results to those already known except for two methods which yielded a product with a markedly superior resistance to thermal degradation. The process according to the present invention relies on a hydrothermal or refluxing treatment of a substantially homogeneous mixture of aluminium and barium containing compounds.

The hydrothermal or refluxing treatment results 15 in crystals of boehmite which appear to be coated with a layer of barium or a barium compound some nanometres thick. The layer is thought to be barium carbonate. Boehmite (AlO.OH) is an aluminium monohydroxide which is a gamma-alumina precursor. On heating to about 500°C, boehmite transforms to gamma-alumina without a 20 substantial change in external crystal shape, but with the development of internal porosity, which is thought to be responsible for the unusually high specific surface area of gamma-alumina. It seems that by 25 depositing the barium as a substantially uniform surface coating, the effectiveness of the barium as a stabilizer for gamma-alumina is maximized and that the hydrothermal or refluxing method is therefore markedly superior to other methods such as calcining that yield a more or less homogeneous mixture of phases. 30

On heating, the hydrothermally treated material transforms to gamma-alumina at temperatures up to about 1000°C (see Figure 3) and transition alumina (theta and delta phases) at temperatures up to about

10

35

1400°C for about 1 hour (see Figure 4). The layer comprising the barium-containing compound, possibly barium carbonate, transforms first to barium oxide on heating to about 830°C in the pure state and then subsequently to barium aluminate.

After heating to a higher temperature, X-ray diffraction showed that the barium formed barium oxide at temperatures up to about 1000°C and various barium aluminates (for example $BaAl_2O_4$, $BaAl_{13.2}O_{20.8}$ and BaAl_{9.2}O_{14.8}) at higher temperatures. These phases were present in all samples, not just in the case of the hydrothermally treated samples.

15 Barium aluminates are believed to be effective in preventing the nucleation of alpha-alumina if a sufficient quantity is present on the surface of gamma-alumina particles to fill surface defects which otherwise act as nucleation sites for the phase 20 transformation to alpha-alumina.

In microwave-treated materials after about 1 hour at approximately 1200°C, barium aluminate (BaAl_{13.2}O_{20.8}) was visible as nano-sized particles predominantly inside much larger alpha-alumina crystals in 25 transmission electron micrographs and was particularly well-developed after about 1 hour at approximately 1400°C (see Figures 9C and 9D), whereas in hydrothermally-treated alumina such particles were absent after treatment at 1400°C, except when heated 30 in argon instead of air, in which case nano-sized particles of barium aluminate (BaAl₁₂O₁₉) were visible, predominantly on the surface of the alpha-alumina crystals (see Figure 2). The conclusion is that barium aluminate was present as a thin surface layer

WO 00/50168 PCT/GB00/00571

- 19 -

in the hydrothermally treated material even after 1 hour at 1400°C in air, but not in a similar starting material treated by oven-drying or microwave-drying because those materials transformed or partly transformed to alpha-alumina after 1 hour at 1200°C, a temperature similar to that reported by other authors. An important step therefore seems to be the formation of a generally uniform surface layer of barium and/or a barium compound on the boehmite crystals during the hydrothermal or refluxing treatment.

The length of hydrothermal treatment is important in determining the specific surface area of the product after the subsequent high temperature 15 treatment. After heating for 1 hour at 1400°C, a material that was previously given a 4 hour hydrothermal treatment at approximately 210°C resulted in theta-alumina with a specific surface area of $6 \text{ m}^2/\text{g}$. In contrast, after heating for 1 hour at 1400°C, a material that was previously given a 22 hour 20 hydrothermal treatment at 210°C resulted in thetaalumina with a more useful specific surface area of 31 m²/g. Such a material would retain sufficient surface area to remain effective as a catalyst support material. Figures 5A and 5B shows that there is 25 remarkably little change in general appearance of the hydrothermally treated material before (5A) and after (5B) heating for 1 hour at 1400°C which demonstrates the effectiveness of this material in resisting 30 thermal degradation.

The present invention will now be described further with reference to the following Examples and Comparative Examples.

5

10

Examples

Aluminium chloride hexahydrate (AlCl₃·6H₂O) (Merck, crystalline, purified), barium chloride dihydrate (BaCl₂·2H₂O) (Baker, analysed, reagent grade), urea (CON₂H₄) (Acros, p.a.) and polyvinylpyrrolidone (PVP, Aldrich, K-30, special grade) were used as received. Solutions of the above reagents were filtered before use.

10

5

Alumina and alumina doped with approximately 10 mol% baria were synthesised by homogeneous precipitation from a metal salt solution induced by the thermal decomposition of urea as shown in the flow chart in Figure 6. On heating the solution to 15 approximately 86°C, the urea decomposed to produce a controlled and uniform increase in pH until precipitation occurred. Very fine sol particles sedimented from the solution after standing for several hours at room temperature. The reaction 20 product was centrifuged to separate the characteristically voluminous and transparent gel. The gel was washed twice with double its own volume of distilled water to remove most of the anion and organic impurities. About 1200 ml of the transparent 25 gel was produced from 5 l of solution in a typical experiment.

Some Ba²⁺ was detected qualitatively in the
washing water by precipitation with SO₄²⁺. The loss of
barium could be avoided by omitting the washing step,
in which case the chloride anion was removed by
sublimation of NH₄Cl, either during microwave drying,
or during calcination in the case of the oven-dried
specimens. For autoclave treated gels, washing

carried out after crystallisation of boehmite and barium carbonate removed chloride ions without a loss of barium.

5 The alumina and alumina-baria gels were treated in a variety of different ways: air-dried at room temperature; oven-dried at a temperature in the range of from 90 to 100°C; dried in a microwave oven; hydrothermally treated in an autoclave. The air-dried 10 gel was prepared by leaving the gel in a glass beaker at room temperature until a solid formed which was then crushed in an agate mortar to produce a fine powder. The oven-dried gel was prepared by heating the gel in air in an oven at about 90 to 100°C for approximately 24 hours; the solid was then crushed as 15 above to produce a fine powder. The microwave-dried samples of gel were prepared in a standard domestic microwave oven at a power setting of 500 W using cycles of 5 min power on and from 2 to 3 min power 20 off, because of extensive foaming of the gel due to the release of gaseous products. The dried solid was crushed as above and later calcined at different temperatures. The hydrothermal treatment was conducted on 150 ml samples of centrifuged gel in 230 25 ml Teflon-lined pressure-tight steel containers. A heating rate of about 0.5°C/min to a temperature of 210 \pm 5°C was used. The hold time at the temperature was about 4 hours. The resulting brownish liquor with a pH in the range of from 8 to 8.5 was decanted and the crystallised product was dried at a temperature 30 from 90 to 100°C. The crystalline product was calcined in either air or argon.

The size of the colloidal particles in the alumina and alumina-baria gels was measured by Small

Angle X-ray Scattering (SAXS); evaluation of the results of this technique was based on the assumption that the gel was a fractal system (see J. Teixeira, J. Appl. Cryst. 21 (1988) 781). The phase composition of the samples was determined by XRD, either in-situ 5 during calcination (Siemens D5000), or conventionally before and after calcination at various temperatures (Philips PW173). Thermal decomposition of the gel to the calcined products on heating to 1400°C was 10 observed by differential thermal analysis and thermogravimetry (DTA/TG, Netsch Simultaneous Analyser STA 409). Transmission electron microscopy (TEM, Philips EM 400) and High Resolution Transmission Electron Microscopy (HRTEM, JEOL 4000 EX/II with a point resolution of 0.165 nm at 400 kV) were used for 15 analysis of the fine structure of uncalcined and calcined samples. Electron microscopy samples were prepared by dispersing the powder in ethyl alcohol, placing a drop of this suspension on a carbon-coated 20 copper grid, and allowing the alcohol to evaporate. The specific surface area was measured by BET nitrogen adsorption (Coulter-Omnisorp 360).

The important characteristic of the homogeneous 25 precipitation process was that it yielded a product substantially uniform both in chemical composition and particle size. The decomposition of urea on heating the solution released ammonium ions into solution in a very uniform manner resulting in a gradual and uniform 30 increase in pH. These uniform conditions ensured that the alumina-baria composite gel which precipitated from the dissolved chloride salts was substantially chemically homogeneous on a colloidal scale. precipitation kinetics were easily controlled through the heating rate and the ageing temperature. 35 The flow

chart in Figure 6 summarises the steps and conditions of the synthesis.

The size of the sol particles in the gel samples was calculated from SAXS measurements as shown in Table 1. The alumina and alumina-baria precursor gels both consisted of particles of about 4 nm diameter and the fractal dimension indicates that both gels were in a similar state of aggregation.

10

5

	Alumina	Alumina- baria
Fractal dimension	2.2	2.4
Particle diameter (nm)	3.7	4.1

15

Table 1: Properties of alumina and alumina-baria precursor gel measured by SAXS

The phase evolution during calcination was studied by X-ray diffraction "in-situ" and conventionally, i.e. on specimens measured at room temperature after calcination. The main results for the conventional measurements are summarised in Table 2 with phases listed in order of abundance. The phases in alumina-baria samples were compared with those in pure alumina samples to obtain information on the effects of each of the gel treatment techniques.

10

25

30

	Phases determined by XRD							
		argon						
Sample	uncalcined	1000°C	1200°C	1400°C	1400°C			
		1h	1h	1h	1h			
oven dried	amorphous	Y-A1203	α-A1,0,	α-A1,0,	nd			
100°C	bayerite	BaO	BaAl ₂ O ₄	β,				
24h	urea (tr)		β,	'	l			
microwave	amorphous	Y-A1203	α-Al ₂ O ₃	α-Al ₂ O ₃	g=310			
dried	-boehmite	β _{II} (tr)	θ-A1 ₂ 0 ₃	β	α-Al ₂ O ₃			
	BaAl ₁₂ O ₁₉ (i)		β,	-1	βι			
autoclaved	boehmite	Y-A1203	0-A1,0,	0-A1,0,	α-Al ₂ O ₃			
4h	BaCO3(i)	δ-Al2O3	δ-Al ₂ O ₃	δ-Al ₂ O ₃	θ-Al ₂ O ₃			
		BaO(i)	BaAl ₂ O ₄	BaAl ₂ O ₄	BaAl ₁₂ O ₁₉			
				β	12 19			
autoclaved	nd	nd	nd	0-A1,0,	nd			
22h	1	j		BaAl,O				
				β,				

 $\beta_{I} = BaA1_{13.2}O_{20.8}, (BaO·6.6Al_{2}O_{3}), \beta_{II} = BaAl_{9.2}O_{14.8},$ $(BaO·4.6Al_{2}O_{3}), tr = traces, (i) only most intense$ peaks visible, nd = not determined

Table 2: Phase composition of alumina-baria samples 20 before and after calcination of dehydrated gel.

The alumina-baria gel dried at room temperature appeared to be amorphous from the XRD spectra. The only indication of the presence of a crystalline phase was a small amount of retained urea. The XRD spectrum for the oven-dried sample (100°C) also indicated predominantly amorphous material and the presence of very poorly crystallized bayerite Al(OH)₃, besides traces of crystalline urea. There was no indication of the presence of any crystalline barium containing

phase.

In the microwave-treated alumina sample, "in situ" XRD indicated the presence of microcrystalline boehmite to approximately 400°C at which temperature 5 the sample became amorphous and remained amorphous to about 800-850°C when transition alumina (Y-Al2O3) appeared. At about 950-1000°C transformation of y- Al_2O_3 to α - Al_2O_3 began and was almost complete at 10 1200°C. In the microwave-treated alumina-baria sample, the phase initially present was also microcrystalline boehmite (or pseudo boehmite) which became amorphous at about 400°C. The transition aluminas that formed were $\gamma-\text{Al}_2\text{O}_3$ and $\theta-\text{Al}_2\text{O}_3$ instead 15 of just γ -Al₂O₃ as in the pure alumina sample. The transformation from $\theta\text{-Al}_2\text{O}_3$ to $\alpha\text{-Al}_2\text{O}_3$ began at about 1050-1100°C. After calcination at 1000°C for 1 hour traces of the BaAl $_{9.2}O_{14.8}$ (β_{11}) phase were present. After calcination for 1 hour at 1200°C or 1400°C, 20 $BaAl_{13.2}O_{20.8}$ (β_1) was present instead of β_{11} . Calcination in argon or argon and air for 1 hour at 1400°C gave the same result.

In the autoclave treated alumina and alumina baria precursor gels XRD showed that the samples
initially consisted of well-crystallised boehmite
which decomposed at 450-500°C to an amorphous phase.
Two rather weak peaks corresponding to the main peaks
in the spectrum for γ-barium carbonate were also
detected. In-situ XRD showed that the two BaCO₃ peaks
persisted to 700°C at which temperature a peak
corresponding to the main BaO peak appeared and
transition alumina formed. After 1 hour at 1000°C, γAl₂O₃, δ-Al₂O₃ and BaO were the phases present. After
1 hour at 1200°C γ-Al₂O₃, δ-Al₂O₃ and BaAl₂O₄ were

present. After 1 hour at $1400 \, ^{\circ}\text{C}$, $\theta-\text{Al}_2\text{O}_3$ and $\delta-\text{Al}_2\text{O}_3$ were present with BaAl_2O_4 and $\text{BaAl}_{9,2}\text{O}_{14.8}$ (β_{11}). A trace amount of $\alpha-\text{Al}_2\text{O}_3$ was found in this sample calcined in air for 1 hour and, after 10 hours at $1400 \, ^{\circ}\text{C}$, $\theta-\text{Al}_2\text{O}_3$ was still the predominant phase. However, in samples calcined in argon for 1 hour at $1400 \, ^{\circ}\text{C}$ the principle phase was $\alpha-\text{Al}_2\text{O}_3$.

The DTA/TG curves in Figure 7 for the hydrothermally treated baria-doped samples show well-10 defined exothermic peaks corresponding to the decomposition of boehmite by dehydroxylation. A displacement of the dehydroxylation peaks to higher temperatures with increased heating rate indicates the slow kinetics of dehydration. The theoretical weight 15 loss for the dehydroxylation of boehmite is 15%. Ba doped boehmite a weight loss of less than 15% was expected whatever barium compound was present, for example carbonate or hydroxide, due to the high atomic weight of barium compared with aluminium. 20 no indication of Ba phases such as BaCO, or Ba(OH)2 XH2O which have a number of well-defined peaks by which they could easily be identified if present.

BET surface area measurements were made on samples calcined for 1 hour at either 1200°C or 1400°C. For calcination at 1200°C, a value of 45.2 m²/g was measured for specimens autoclaved for 4 hours. For calcination at 1400°C, values of 5.9 m²/g and 31.2 m²/g were obtained for specimens autoclaved for 4 hours and 22 hours respectively. The longer autoclave treatment resulted in larger boehmite crystal size in the sample before calcination. After calcination at 1400°C, the phase composition of the specimens was also different. From XRD analysis, the

phases present in order of abundance in the specimen autoclaved for 4h were: θ -Al₂O₃, δ -Al₂O₃, BaAl₂O₄ and β ₁₁. The specimen autoclaved for 22 hours differed in that δ -Al₂O₃ was absent.

5

TEM examination showed that considerable morphological differences existed between oven-dried, microwave-dried and hydrothermally treated samples, both before and after calcination.

10

15

20

Before calcination, the oven-dried material consisted of densely agglomerated fine crystals, Figures 8A and 8B, which were identified by electron diffraction as bayerite. The microwaved material was much less tightly agglomerated as shown in Figure 9A and consisted of boehmite crystallites of about 5 nm as seen by HRTEM (not shown here) which approximately correspond to the size of the colloidal particles in the gel. The hydrothermally treated sample consisted of plate-like boehmite crystals about 250 x 100 x 10 nm as shown in Figure 10A. There was no evidence of a separate Ba containing phase in any of the uncalcined samples.

25

30

35

TEM micrographs of oven-dried material calcined for 1 hour at $1000\,^{\circ}$ C indicated the presence of a γ - Al_2O_3 matrix and finely distributed BaO (small dark particles) (see Figures 8C and 8D). The crystallite size of the γ - Al_2O_3 and barium containing phase was about the same after calcination as that of bayerite before calcination (see Figures 8B and 8D). After calcination at $1000\,^{\circ}$ C the autoclaved material consisted of γ - Al_2O_3 which retained the shape of the boehmite crystals (pseudomorphs) as shown in Figure 10B. The pseudomorphs were porous and the pore facets

can be seen in Figure 10C.

The TEM images of microwaved material calcined for 1 hour at 1200°C, Figures 1 and 9B, show the start of the transition from $\theta-\text{Al}_2\text{O}_3$ to $\alpha-\text{Al}_2\text{O}_3$ in pure 5 alumina and in alumina-baria samples, respectively. After calcination at 1400°C for 1 hour in air or argon, the microwaved material consisted of BaAl_{13.2}O_{20.8} (β_{1}) particles of from 10 to 15 nm diameter and much larger α -Al₂O₃ crystals as shown in Figures 9C to 9F. 10 The β_1 particles were predominantly within the α -Al₂O₃ crystals after calcination in air, but were distributed both within and on the surface of the α -Al₂O₃ crystals after calcination in argon.

15

35

The autoclave treated material, calcined at 1400°C in air, consisted mainly of finely twinned θ -Al203 with twin planes spaced about 5 nm apart as shown in Figures 11 A-D. The boehmite pseudomorph crystal shape was still retained although the internal 20 porosity was much less than at 1000°C, and was more spherical in shape. The θ -Al,0, twin planes were parallel to the long axis of the pseudomorphs, i.e. the boehmite <001> direction. There was no evidence of sintering between the crystals and their size was 25 similar to that at 1200°C. Small amounts α -Al,0, were observed. Detailed HRTEM examination of transition alumina structures in the autoclave treated samples indicated that the fine twinning in 0-Al203 seen in the 30 material calcined at 1400°C probably resulted from the ordering of cation vacancies in 6-Al,0, at lower temperatures. In contrast, autoclave treated material, calcined at 1400°C in argon, consisted of large crystals of α -Al₂O₃ decorated on the surface with 10 nm particles of $BaA1_{12}O_{10}$ which has a

magnetoplumbite structure rather than a β type structure (see Figures 12A and 12B).

The present invention provides a process whereby transition alumina can be further stabilized against thermal degradation to alpha-alumina. This results in a catalyst which is effective at higher operating temperatures and for longer times compared with the prior art catalysts. An additional advantage of the present invention is that it is very effective at forming thin (approximately 1 to 5 μ m) washcoats of gamma-alumina on cordierite or other ceramic substrates. This is important because the use in the future of ceramic honeycomb substrates with a very high channel density will require the use of thinner washcoats. The process according to the present invention can produce thinner films where (coated) boehmite is used in the slurry, rather than gammaalumina.

20

5

10

15

25

30

Claims:

5

10

15

- 1. A process for the production of a precursor material for use in the preparation of a stabilized alumina catalyst support material, which process comprises the steps of:
- (a) providing a mixture comprising aluminium and barium containing compounds; and
- (b) heating the mixture under conditions to form particles of boehmite at least partially coated with a layer comprising barium and/or a compound of barium.
- 2. A process as claimed in claim 1, wherein the mixture comprising aluminium and barium containing compounds is a substantially homogenous mixture.
- 3. A process as claimed in claim 1 or claim 2, wherein the layer formed on the particles of boehmite comprises barium carbonate.
- 4. A process as claimed in any one of the preceding claims, wherein the aluminium containing compound comprises aluminium chloride, aluminium nitrate, aluminium sulphate, alumina and/or boehmite.
- 5. A process as claimed in any one of the preceding claims, wherein the barium containing compound comprises barium chloride, barium sulphate and/or barium nitrate.
- A process as claimed in any one of the preceding
 claims, wherein the mixture comprising aluminium and

barium containing compounds is in the form of a substantially homogeneous precipitate.

- 7. A process as claimed in claim 6, wherein the substantially homogeneous precipitate is formed by a process involving homogeneous precipitation of aluminium and barium containing compounds from salt solution.
- 8. A process as claimed in claim 7, wherein the homogeneous precipitation involves generating a base within the salt solution through thermal decomposition of a water-soluble reagent.
- 9. A process as claimed in claim 8, wherein the water-soluble reagent comprises urea and/or hexamethylene tetramine.
- 10. A process as claimed in any one of claims 7 to 9,
 wherein the salt solution further includes a
 dispersant, such as polyvinylpyrrolidone.
- 11. A process as claimed in any one of claims 1 to 5, wherein the mixture in step (a) is formed by adding a solution comprising oxalic acid and a water soluble barium compound, preferably barium nitrate, to an aqueous slurry comprising boehmite.
- 12. A process as claimed in any one of the preceding claims, wherein heating of the mixture in step (b) is achieved by a hydrothermal process.
- 13. A process as claimed in claim 12, wherein the hydrothermal treatment is carried out at a temperature
 35 in the range of from 90 to 300°C, more preferably from

25

30

35

100 to 220°C.

- 14. A process as claimed in claim 12 or claim 13, wherein the hydrothermal treatment is carried out at a pressure in the range of from 1 to 150 bar, preferably from 5 to 50 bar.
- 15. A process as claimed in any one of claim 12 to 14, wherein the hydrothermal treatment is carried out 10 for a time in the range of from 30 minutes to 25 hours, preferably from 1 to 10 hours.
- 16. A process as claimed in any one of claims 1 to 11, wherein the mixture in step (b) is heated at a temperature in the range of from ambient temperature to boiling temperature, preferably by a reflux process.
- 17. A process for the preparation of a stabilized alumina catalyst support material, which process comprises the steps of:
 - (i) providing a precursor material comprising particles of boehmite at least partially coated with a layer comprising barium and/or a compound of barium; and
 - (ii) heating the coated particles of boehmite to a temperature at which at least some of the boehmite transforms to gamma-alumina.
 - 18. A process as claimed in claim 17, comprising the step of further heating whereby at least some of the gamma-alumina transforms to theta and/or delta-alumina.

- 19. A process as claimed in claim 17 or claim 18, wherein said precursor material is produced according to a process as claimed in any one of claim 1 to 16.
- 20. A process for the preparation of an automotive catalytic converter, which process comprises the steps of dispersing a mixture comprising particles of boehmite at least partially coated with a layer comprising barium and/or a compound of barium on a
- metal or ceramic monolithic substrate, followed by heating the precursor mixture to a temperature at which at least some of the boehmite transforms to gamma-alumina.
- 21. A process as claimed in claim 20, comprising the step of further heating whereby at least some of the gamma-alumina transforms to theta and/or delta-alumina.
- 20 22. A process as claimed in claim 17 or claim 20, wherein heating is carried out at a temperature in the range of from 500 to 1000°C.
- 23. A process as claimed in claim 18 or claim 21, wherein the further heating is carried out at a temperature in the range of from 1000 to 1400°C.
- 24. A process for the preparation of a catalyst, such as an automotive catalytic converter, which process comprises the step of dispersing a stabilized alumina catalyst support material prepared according to the process of any one of claims 17 to 19 on a metal or ceramic monolithic substrate.
- 35 25. A precursor material for use in the preparation

of catalyst support material, which precursor material comprises particles of boehmite and/or transition alumina at least partially coated with a layer comprising barium and/or a compound of barium.

5

26. A precursor material as claimed in claim 25, wherein the layer formed on the particles of boehmite and/or transition alumina comprises barium carbonate, barium oxide and/or barium aluminate.

10

- 27. A precursor material as claimed in claim 25 or claim 26 provided in the form of a slurry or washcoat.
- 28. A stabilized alumina catalyst support material
 which comprises gamma, theta and/or delta-alumina, and
 which withstands substantial degradation to alphaalumina after exposure to a temperature of
 approximately 1400°C for 1 hour.
- 29. A stabilized alumina catalyst support material as claimed in claim 28, wherein particles of gamma, theta and/or delta-alumina are at least partially coated with a layer comprising barium carbonate, barium oxide and/or barium aluminate.

25

30. A stabilized alumina catalyst carrier or support material as claimed in claim 28 or claim 29 having a specific surface area of at least 20 m²/g after exposure to a temperature of 1400°C for 1 hour.

30

31. A stabilized alumina catalyst carrier or support material as claimed in any one of claims 28 to 30 having a specific surface area of at least 31 $\rm m^2/g$ after exposure to a temperature of 1400°C for 1 hour.

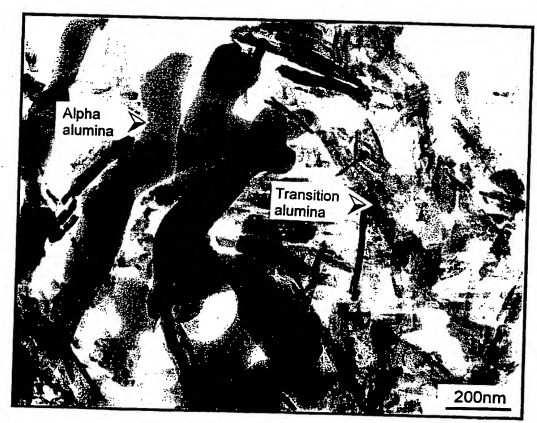


FIG. 1

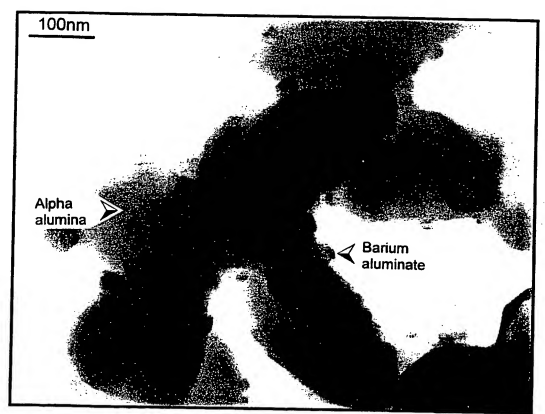


FIG. 2

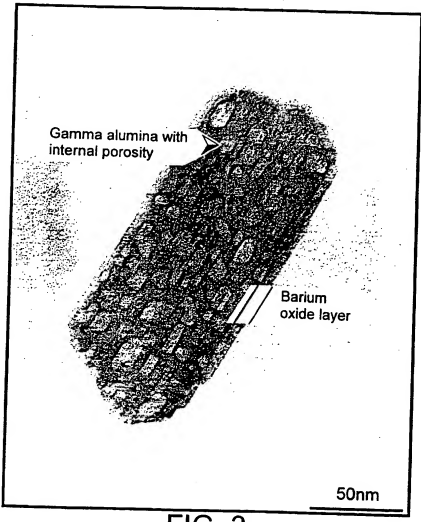


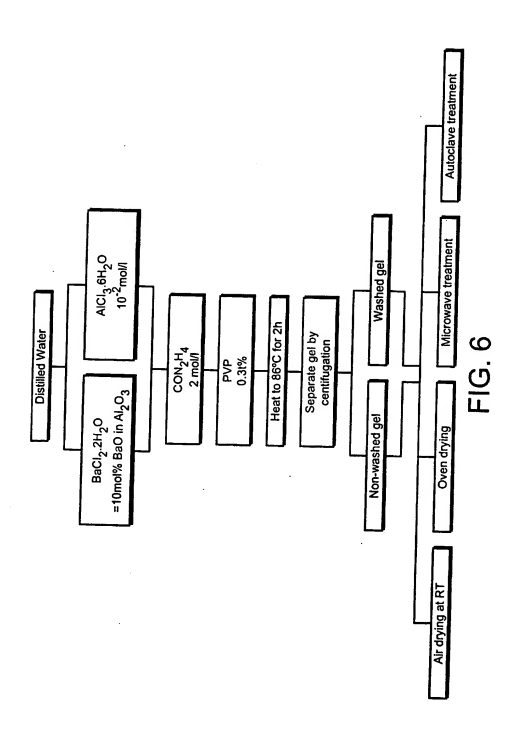
FIG. 3

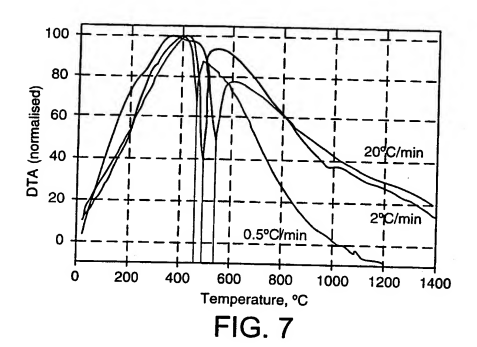


SUBSTITUTE SHEET (RULE 26)

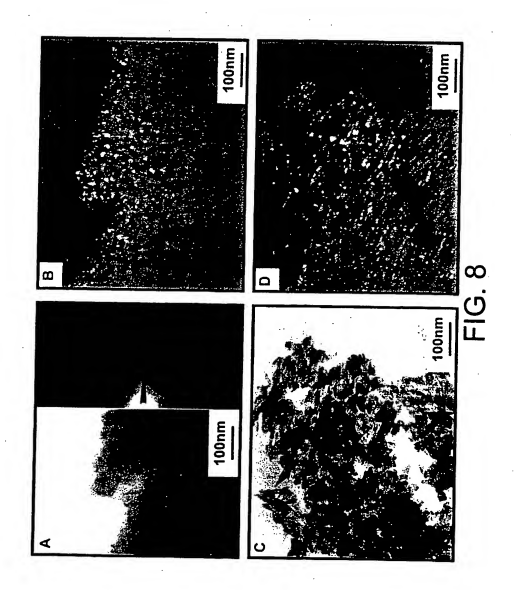


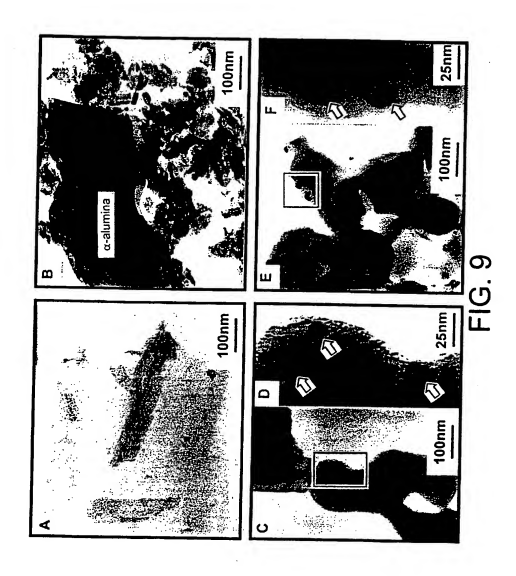
SUBSTITUTE SHEET (RULE 26)



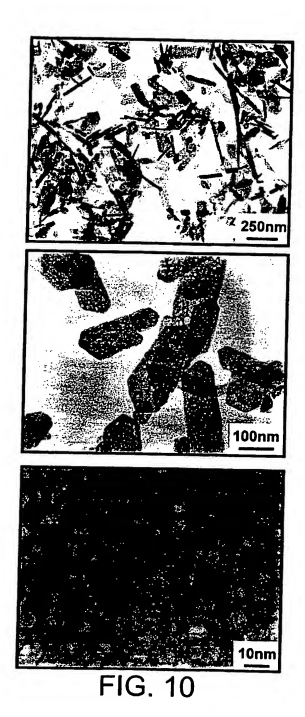


SUBSTITUTE SHEET (RULE 26)

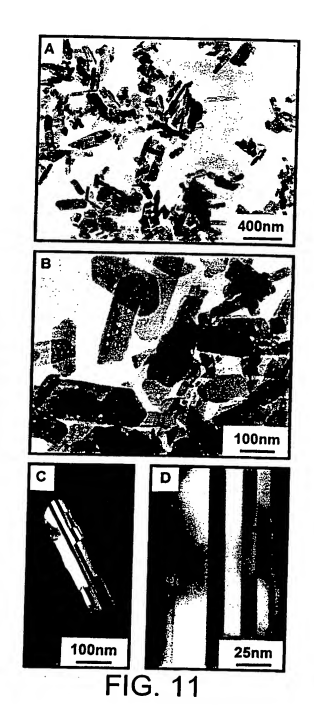




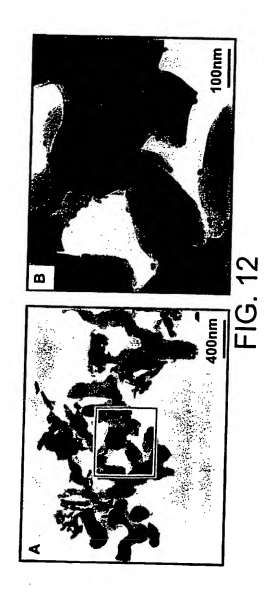
SUBSTITUTE-SHEET (RULE 26)



SUBSTITUTE SHEET (RULE 26)



SUBSTITUTE SHEET (RULE 26)



INTERNATIONAL SEARCH REPORT

Inten nal Application No PCT/GB 00/00571

			101/46 0	07 00 5 7 1							
A CLASS IPC 7	FIGATION OF SUBJECT MATTER B01J23/02 B01J23/58 B01J37	7/02 B01D	53/94								
According to International Patent Classification (IPC) or to both national classification and IPC											
B. FIELDS SEARCHED											
IPC 7	ocumentation searched (classification system followed by classific B01J										
	tion searched other than minimum documentation to the extent the										
WPI Da	ata base consulted during the international search (name of data	base and, where pra	ctical, search terms used								
C. DOCUM	ENTS CONSIDERED TO BE RELEVANT										
Category *	Citation of document, with indication, where appropriate, of the	relevant passages		Relevant to claim No.							
X	DATABASE WPI Section Ch, Week 199040 Derwent Publications Ltd., Londo Class H06, AN 1990-302272 XP002140068 & JP 02 214541 A (TOYO KOGYO CO 27 August 1990 (1990-08-27)			1,2,17, 19,20, 24,27							
Y	abstract			4-6							
Υ	US 5 212 130 A (ADDIEGO WILLIAM 18 May 1993 (1993-05-18) claim 1	P ET AL)		4-6							
A	US 5 116 800 A (SUMMERS II JACK 26 May 1992 (1992-05-26)	C ET AL)									
Further documents are listed in the continuation of box C. Patent family members are listed in annex.											
"A" document conside "E" earlier do filing dat "L" document which is citation of document other me	which may throw doubts on priority claim(s) or cited to establish the publication date of another or other special reason (as specified) or referring to an oral disclosure, use, exhibition or sans	T' later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention X' document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone occurrent of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art.									
later tha	n the priority date claimed	*& * document memi	per of the same patent for	amily							
	June 2000	Date of mailing of the international search report 28/06/2000									
Name ema	lling address of the ISA European Patent Office, P.B. 5818 Patentlaan 2 NL – 2280 HV Rijswijk	Authorized officer									
	Tel. (+31-70) 340-2040, Tx. 31 651 epo ni, Fax: (+31-70) 340-3016	Thion	M								

INTERNATIONAL SEARCH REPORT

information on patent family members

inter mai Application No PCT/GB 00/00571

Patent document cited in search report		Publication date	Patent family member(s)		Publication date
JP 2214541	A	27-08-1990	JP	2744275 B	28-04-1998
US 5212130	A	18-05-1993	EP JP	0560074 A 6000372 A	15-09-1993 11-01-1994
US 5116800	A	26-05-1992	DE DE EP MX WO	69104062 D 69104062 T 0561904 A 9102196 A 9210271 A	20-10-1994 23-03-1995 29-09-1993 01-06-1992 25-06-1992

This Page is Inserted by IFW Indexing and Scanning Operations and is not part of the Official Record

BEST AVAILABLE IMAGES

Defective images within this document are accurate representations of the original documents submitted by the applicant.

Defects in the images include but are not limited to the items checked:

| BLACK BORDERS
| IMAGE CUT OFF AT TOP, BOTTOM OR SIDES
| FADED TEXT OR DRAWING
| BLURRED OR ILLEGIBLE TEXT OR DRAWING
| SKEWED/SLANTED IMAGES
| COLOR OR BLACK AND WHITE PHOTOGRAPHS
| GRAY SCALE DOCUMENTS
| LINES OR MARKS ON ORIGINAL DOCUMENT
| REFERENCE(S) OR EXHIBIT(S) SUBMITTED ARE POOR QUALITY

IMAGES ARE BEST AVAILABLE COPY.

OTHER:

As rescanning these documents will not correct the image problems checked, please do not report these problems to the IFW Image Problem Mailbox.